

# **Dy-Mark**

Chemwatch Hazard Alert Code: 4

Chemwatch: 42-9966 Version No: 11.1

Safety Data Sheet according to WHS Regulations (Hazardous Chemicals) Amendment 2020 and ADG requirements

Issue Date: 20/04/2022 Print Date: 20/04/2022 S.GHS.AUS.EN.E

## SECTION 1 Identification of the substance / mixture and of the company / undertaking

#### **Product Identifier**

Product name	Dy-Mark Protech Chain Lubricant
Chemical Name	Not Applicable
Synonyms	Product Code: 42033003
Proper shipping name	AEROSOLS
Chemical formula	Not Applicable
Other means of identification	Not Available

#### Relevant identified uses of the substance or mixture and uses advised against

Relevant identified uses	Application is by spray atomisation from a hand held aerosol pack
	Use according to manufacturer's directions.

## Details of the supplier of the safety data sheet

Registered company name	Dy-Mark
Address	89 Formation Street Wacol QLD 4076 Australia
Telephone	+61 7 3327 3004
Fax	+61 7 3327 3009
Website	http://www.dymark.com.au
Email	info@dymark.com.au

## Emergency telephone number

Association / Organisation	Dy-Mark
Emergency telephone numbers	+61 7 3327 3099
Other emergency telephone numbers	Not Available

#### **SECTION 2 Hazards identification**

#### Classification of the substance or mixture

## HAZARDOUS CHEMICAL. DANGEROUS GOODS. According to the WHS Regulations and the ADG Code.

#### ChemWatch Hazard Ratings

	Min	Max	
Flammability	4		
Toxicity	1	1	0 = Minimum
Body Contact	2	1	1 = Low
Reactivity	2	1	2 = Moderate
Chronic	1		3 = High 4 = Extreme

Poisons Schedule	Not Applicable
Classification <sup>[1]</sup>	Aerosols Category 1, Aspiration Hazard Category 1, Skin Corrosion/Irritation Category 2, Specific Target Organ Toxicity - Single Exposure (Narcotic Effects) Category 3, Hazardous to the Aquatic Environment Long-Term Hazard Category 2
Legend:	1. Classified by Chernwatch; 2. Classification drawn from HCIS; 3. Classification drawn from Regulation (EU) No 1272/2008 - Annex VI

#### Label elements



Signal word Danger

## Hazard statement(s)

• •	
AUH044	Risk of explosion if heated under confinement.
H222+H229	Extremely flammable aerosol. Pressurized container: may burst if heated.
H304	May be fatal if swallowed and enters airways.
H315	Causes skin irritation.
H336	May cause drowsiness or dizziness.
H411	Toxic to aquatic life with long lasting effects.

#### Precautionary statement(s) Prevention

P210	Keep away from heat, hot surfaces, sparks, open flames and other ignition sources. No smoking.
P211	Do not spray on an open flame or other ignition source.
P251	Do not pierce or burn, even after use.
P271	Use only outdoors or in a well-ventilated area.
P261	Avoid breathing mist/vapours/spray.
P273	Avoid release to the environment.
P280	Wear protective gloves and protective clothing.
P264	Wash all exposed external body areas thoroughly after handling.

#### Precautionary statement(s) Response

P301+P310	IF SWALLOWED: Immediately call a POISON CENTER/doctor/physician/first aider.	
P331	Do NOT induce vomiting.	
P312	Call a POISON CENTER/doctor/physician/first aider/if you feel unwell.	
P391	Collect spillage.	
P302+P352	IF ON SKIN: Wash with plenty of water and soap.	
P304+P340	4+P340 IF INHALED: Remove person to fresh air and keep comfortable for breathing.	
P332+P313	If skin irritation occurs: Get medical advice/attention.	
P362+P364	Take off contaminated clothing and wash it before reuse.	

#### Precautionary statement(s) Storage

P405	Store locked up.
P410+P412	Protect from sunlight. Do not expose to temperatures exceeding 50 °C/122 °F.
P403+P233	Store in a well-ventilated place. Keep container tightly closed.

#### Precautionary statement(s) Disposal

P501

Dispose of contents/container to authorised hazardous or special waste collection point in accordance with any local regulation.

# **SECTION 3 Composition / information on ingredients**

## Substances

See section below for composition of Mixtures

# Mixtures

CAS No	%[weight]	Name
8042-47-5	10-15	white mineral oil (petroleum)
Not Available	20-30	poly alpha olefin
107-83-5	20-30	2-methylpentane
68476-85-7.	20-35	LPG (liquefied petroleum gas)
Legend:	1. Classified by Chemwatch; 2. Classification drawn from HCIS; 3. Classification drawn from Regulation (EU) No 1272/2008 - Annex VI; 4. Classification drawn from C&L * EU IOELVs available	

# **SECTION 4 First aid measures**

Description of first aid measures		
Eye Contact	<ul> <li>If aerosols come in contact with the eyes:</li> <li>Immediately hold the eyelids apart and flush the eye continuously for at least 15 minutes with fresh running water.</li> <li>Ensure complete irrigation of the eye by keeping eyelids apart and away from eye and moving the eyelids by occasionally lifting the upper and lower lids.</li> </ul>	

	<ul> <li>Transport to hospital or doctor without delay.</li> <li>Removal of contact lenses after an eye injury should only be undertaken by skilled personnel.</li> </ul>
Skin Contact	If solids or aerosol mists are deposited upon the skin: <ul> <li>Flush skin and hair with running water (and soap if available).</li> <li>Remove any adhering solids with industrial skin cleansing cream.</li> <li>DO NOT use solvents.</li> <li>Seek medical attention in the event of irritation.</li> </ul>
Inhalation	<ul> <li>If aerosols, fumes or combustion products are inhaled:</li> <li>Remove to fresh air.</li> <li>Lay patient down. Keep warm and rested.</li> <li>Prostheses such as false teeth, which may block airway, should be removed, where possible, prior to initiating first aid procedures.</li> <li>If breathing is shallow or has stopped, ensure clear airway and apply resuscitation, preferably with a demand valve resuscitator, bag-valve mask device, or pocket mask as trained. Perform CPR if necessary.</li> <li>Transport to hospital, or doctor.</li> </ul>
Ingestion	<ul> <li>Avoid giving milk or oils.</li> <li>Avoid giving alcohol.</li> <li>Not considered a normal route of entry.</li> <li>If spontaneous vomiting appears imminent or occurs, hold patient's head down, lower than their hips to help avoid possible aspiration of vomitus.</li> </ul>

#### Indication of any immediate medical attention and special treatment needed

Treat symptomatically.

+ Heavy and persistent skin contamination over many years may lead to dysplastic changes. Pre-existing skin disorders may be aggravated by exposure to this product.

In general, emesis induction is unnecessary with high viscosity, low volatility products, i.e. most oils and greases.

High pressure accidental injection through the skin should be assessed for possible incision, irrigation and/or debridement.

**NOTE:** Injuries may not seem serious at first, but within a few hours tissue may become swollen, discoloured and extremely painful with extensive subcutaneous necrosis. Product may be forced through considerable distances along tissue planes.

#### **SECTION 5 Firefighting measures**

#### Extinguishing media

SMALL FIRE:

Water spray, dry chemical or CO2

LARGE FIRE:

Water spray or fog.

#### Special hazards arising from the substrate or mixture

**Fire Fighting** 

Fire Incompatibility	+ Avoid contamination with oxidising agents i.e. nitrates, oxidising acids, chlorine bleaches, pool chlorine etc. as ignition may result
dvice for firefighters	
	Alert Fire Brigade and tell them location and nature of hazard.

May be violently or explosively reactive.	
---	--

- Wear breathing apparatus plus protective gloves.
  - Prevent, by any means available, spillage from entering drains or water course.
  - ▶ If safe, switch off electrical equipment until vapour fire hazard removed.
  - Use water delivered as a fine spray to control fire and cool adjacent area.
  - **DO NOT** approach containers suspected to be hot.
  - Cool fire exposed containers with water spray from a protected location.
  - If safe to do so, remove containers from path of fire.
  - Equipment should be thoroughly decontaminated after use.

GENERAL

- Alert Fire Brigade and tell them location and nature of hazard.
- May be violently or explosively reactive.
- Wear breathing apparatus plus protective gloves.
- Consider evacuation
- Fight fire from a safe distance, with adequate cover.
- If safe, switch off electrical equipment until vapour fire hazard removed.
- Use water delivered as a fine spray to control fire and cool adjacent area.
- DO NOT approach cylinders suspected to be hot.
- Cool fire-exposed cylinders with water spray from a protected location.
- If safe to do so, remove containers from path of fire.

FIRE FIGHTING PROCEDURES:

- -----
  - The only safe way to extinguish a flammable gas fire is to stop the flow of gas.
  - If the flow cannot be stopped, allow the entire contents of the cylinder to burn while cooling the cylinder and surroundings with water from a suitable distance.
  - Extinguishing the fire without stopping the gas flow may permit the formation of ignitable or explosive mixtures with air. These mixtures may propagate to a source of ignition.

SPECIAL HAZARDS

- ▶ Excessive pressures may develop in a gas cylinder exposed in a fire; this may result in explosion.
- Cylinders with pressure relief devices may release their contents as a result of fire and the released gas may constitute a further source of hazard for the fire-fighter.
- Cylinders without pressure-relief valves have no provision for controlled release and are therefore more likely to explode if exposed to fire.

FIRE FIGHTING REQUIREMENTS:

	The need for proximity, entry and flash-over protection and special protective clothing should be determined for each incident, by a competent fire-fighting safety professional.
Fire/Explosion Hazard	<ul> <li>Liquid and vapour are highly flammable.</li> <li>Severe fire hazard when exposed to heat or flame.</li> <li>Vapour forms an explosive mixture with air.</li> <li>Severe explosion hazard, in the form of vapour, when exposed to flame or spark.</li> <li>Vapour may travel a considerable distance to source of ignition.</li> <li>Heating may cause expansion or decomposition with violent container rupture.</li> <li>Aerosol cans may explode on exposure to naked flames.</li> <li>Rupturing containers may rocket and scatter burning materials.</li> <li>Hazards may not be restricted to pressure effects.</li> <li>May emit acrid, poisonous or corrosive fumes.</li> <li>On combustion, may emit toxic fumes of carbon monoxide (CO).</li> <li>Combustion products include:</li> <li>carbon dioxide (CO2)</li> <li>other pyrolysis products typical of burning organic material.</li> <li>Contains low boiling substance: Closed containers may rupture due to pressure buildup under fire conditions.</li> <li>May emit clouds of acrid smoke</li> <li>CARE: Water in contact with hot liquid may cause foaming and a steam explosion with wide scattering of hot oil and possible severe burns.</li> <li>Foaming may cause overflow of containers and may result in possible fire.</li> </ul>
HAZCHEM	Not Applicable

# **SECTION 6** Accidental release measures

# Personal precautions, protective equipment and emergency procedures

See section 8

## **Environmental precautions**

See section 12

## Methods and material for containment and cleaning up

Minor Spills	<ul> <li>Clean up all spills immediately.</li> <li>Avoid breathing vapours and contact with skin and eyes.</li> <li>Wear protective clothing, impervious gloves and safety glasses.</li> <li>Shut off all possible sources of ignition and increase ventilation.</li> <li>Wipe up.</li> <li>If safe, damaged cans should be placed in a container outdoors, away from all ignition sources, until pressure has dissipated.</li> <li>Undamaged cans should be gathered and stowed safely.</li> </ul>
Major Spills	<ul> <li>Do NOT exert excessive pressure on valve; Do NOT attempt to operate damaged valve.</li> <li>Clear area of personnel and move upwind.</li> <li>Alert Fire Brigade and tell them location and nature of hazard.</li> <li>May be violently or explosively reactive.</li> <li>Wear breathing apparatus plus protective gloves.</li> <li>Prevent, by any means available, spillage from entering drains or water courses</li> <li>No smoking, naked lights or ignition sources.</li> <li>Increase ventilation.</li> <li>Stop leak if safe to do so.</li> <li>Water spray or fog may be used to disperse / absorb vapour.</li> <li>Absorb or cover spill with sand, earth, inert materials or vermiculite.</li> <li>If safe, damaged cans should be placed in a container outdoors, away from ignition sources, until pressure has dissipated.</li> <li>Undamaged cans should be gathered and stowed safely.</li> <li>Collect residues and seal in labelled drums for disposal.</li> <li>Clear area of all unprotected personnel and move upwind.</li> <li>Alert Emergency Authority and advise them of the location and nature of hazard.</li> <li>May be violently or explosively reactive.</li> <li>Wear full body clothing with breathing apparatus.</li> <li>Prevent by any means available, spillage from entering drains and water-courses.</li> <li>Consider evacuation.</li> <li>Shut off all possible sources of ignition and increase ventilation.</li> <li>No smoking or naked lights within area.</li> <li>Use extreme caution to prevent violent reaction.</li> <li>Stop leak only if safe to so do.</li> <li>Water spray or fog may be used to disperse vapour.</li> <li>Do NOT enter confined space where gas may have collected.</li> <li>Keep area clear until gas has dispersed.</li> <li>Release pressure under safe, controlled conditions by opening the valve.</li> </ul>

Personal Protective Equipment advice is contained in Section 8 of the SDS.

# SECTION 7 Handling and storage

Precautions for safe handling	
Safe handling	The conductivity of this material may make it a static accumulator., A liquid is typically considered nonconductive if its conductivity is below 100 pS/m and is considered semi-conductive if its conductivity is below 10 000 pS/m., Whether a liquid is nonconductive or semi-conductive, the precautions are the same., A number of factors, for example liquid temperature, presence of contaminants, and anti-static additives can greatly influence the conductivity of a liquid. <ul> <li>Avoid all personal contact, including inhalation.</li> <li>Wear protective clothing when risk of exposure occurs.</li> <li>Use in a well-ventilated area.</li> <li>Prevent concentration in hollows and sumps.</li> </ul>

	DO NOT enter confined spaces until atmosphere has been checked.
	Avoid smoking, naked lights or ignition sources.
	Avoid contact with incompatible materials.
	When handling, <b>DO NOT</b> eat, drink or smoke.
	DO NOT incinerate or puncture aerosol cans.
	DO NOT spray directly on humans, exposed food or food utensils.
	Avoid physical damage to containers.
	Always wash hands with soap and water after handling.
	Work clothes should be laundered separately.
	Use good occupational work practice.
	Observe manufacturer's storage and handling recommendations contained within this SDS.
	Atmosphere should be regularly checked against established exposure standards to ensure safe working conditions are maintained.
	Keep dry to avoid corrosion of cans. Corrosion may result in container perforation and internal pressure may eject contents of can
	Store in original containers in approved flammable liquid storage area.
	DO NOT store in pits, depressions, basements or areas where vapours may be trapped.
	No smoking, naked lights, heat or ignition sources.
	Keep containers securely sealed. Contents under pressure.
Other information	Store away from incompatible materials.
Other Information	Store in a cool, dry, well ventilated area.
	Avoid storage at temperatures higher than 40 deg C.
	Store in an upright position.
	Protect containers against physical damage.
	Check regularly for spills and leaks.
	Observe manufacturer's storage and handling recommendations contained within this SDS.

#### Conditions for safe storage, including any incompatibilities

	Suitable contain	er ► Aer ► Che	osol dispenser. eck that containe	ers are clearly la	belled.		
Stora	age incompatibili	y · CARE material · Oil lea about 44 · Autoig • Avo	: Water in conta . Resultant over ks in a pressuriz 5 g/m3 nition temperatu id reaction with	ct with heated m flow of contained red circuit may re rres may be sign oxidising agents	aterial may cau rs may result in esult in a fine fla ificantly lower u	se foaming or a stea fire. Immable spray (the nder particular conc	am explosion with possible severe burns from wide scattering of hot lower flammability limit for oil mist is reached for a concentration of litions (slow oxidation on finely divided materials
$\wedge$	$\wedge$					$\wedge$	



X — Must not be stored together

0 — May be stored together with specific preventions

+ - May be stored together

Note: Depending on other risk factors, compatibility assessment based on the table above may not be relevant to storage situations, particularly where large volumes of dangerous goods are stored and handled. Reference should be made to the Safety Data Sheets for each substance or article and risks assessed accordingly.

## **SECTION 8 Exposure controls / personal protection**

#### **Control parameters**

#### Occupational Exposure Limits (OEL)

#### INGREDIENT DATA

Source	Ingredient	Material name	TWA	STEL	Peak	Notes
Australia Exposure Standards	white mineral oil (petroleum)	Oil mist, refined mineral	5 mg/m3	Not Available	Not Available	Not Available
Australia Exposure Standards	2-methylpentane	Hexane, other isomers	500 ppm / 1760 mg/m3	3500 mg/m3 / 1000 ppm	Not Available	Not Available
Australia Exposure Standards	LPG (liquefied petroleum gas)	LPG (liquified petroleum gas)	1000 ppm / 1800 mg/m3	Not Available	Not Available	Not Available

Emergency Limits

Ingredient	TEEL-1	TEEL-2		TEEL-3
white mineral oil (petroleum)	140 mg/m3	1,500 mg/m3		8,900 mg/m3
2-methylpentane	1,000 ppm	11000** ppm		66000*** ppm
LPG (liquefied petroleum gas)	65,000 ppm	2.30E+05 ppm		4.00E+05 ppm
Ingredient	Original IDLH		Revised IDLH	
white mineral oil (petroleum)	2,500 mg/m3		Not Available	
2-methylpentane	Not Available		Not Available	
LPG (liquefied petroleum gas)	2,000 ppm		Not Available	

#### Exposure controls

Appropriate engineering controls CARE: Use of a quantity of this material in confined space or poorly ventilated area, where rapid build up of concentrated atmosphere may occur, could require increased ventilation and/or protective gear

Engineering controls are used to remove a hazard or place a barrier between the worker and the hazard. Well-designed engineering controls can be highly effective in protecting workers and will typically be independent of worker interactions to provide this high level of protection.

"addr and remover" are in the work environment. Verification can remove or duits an air containment of designed property. The design of a distribution hyper mutanch the particular process and definition or containment in the air containment. We can be address of contains the intervence of address of a distribution is wereboute or doined storage areas. An approved respirator. Correct fit is essential dobian degrade address or distribution is wereboute or doined storage areas. An containment generated in the workplute processors wying "escope" velocities which, in turn, determine the "capture velocities" of treph containment.         Type of Containment:       Speed:         accords, forbased at two velocity into zero of active generation into zero of applies which, in turn, determine the "capture velocities" of treph containment.         Type of Containment:       Speed:         accords, forbased and two velocity into zero of active generation into zero of applies at motion       0.64 min.         Accords relations of two velocity into zero of active generation into zero of applies at motion       1.25 mis (200 500 fmi).         With each range the appropriate value depends on:       2. Contaminante of two velocity were of nucleon velocity or 1.25 mis (200 500 fmi).         With each range the appropriate value depends on:       3. High production. Have and two servers and the containments of high toxicity		The basic types of engineering controls are: Process controls which involve changing the way a job activi Enclosure and/or isolation of emission source which keeps a	ty or process is done to reduce the risk. selected hazard "physically" away from the worker	and ventilation that strategically			
General enhands is adequate under normal conditions. If this of overexposure exists, wear SAA approved respirator. Correct II is essential in its initian indequipe projection.       Provide adequate ventilation in weakhouse or closed storage areas.         Novide adequate ventilation in weakhouse or closed storage areas.       Senter intervention in the unit projection intervention or active texts of ventilation in the unit projection intervention of the conduminant.       Senter         Type of Constminants:       Senter       Senter         accords, (released at low velocity into zone of active generation into zone of rapid at motion)       1-2.5 m/s (200-500 fml Vinto)         1. Boom in currents minimal or towardade to capute.       Upper and of the range       Upper and of the range         1. Room in currents minimal or towardade to capute.       2. Contaminants of the obsolty of or Indivationan ovalue or dy.       2. Contaminants of the obsolty of on analysis of the obsolty of on analysis of the obsolty of on analysis of diatance from the extraction pair (in single cases). Therefore the air special in the obsolt is on analysis of diatance from the extraction pair (in single cases). Therefore the air special in the obsolt is on analysis of diatance from the extraction pair (in single cases). Therefore the air special in the extraction pair (in single cases). Therefore the air special in the obsolt of on analysis of diatance from the extraction pair (in single cases). Therefore the air special in the extraction pair (in single cases). Therefore the air special in the extraction pair (in single cases). Therefore the air special in the extraction pair (in single cases). Therefore the air special in the extraction pair (in single cases). Therefore the air special		"adds" and "removes" air in the work environment. Ventilation can remove or dilute an air contaminant if designed properly. The design of a ventilation system must match the particular process and chemical or contaminant in use. Employers may need to use multiple types of controls to prevent employee overexposure.					
Extra time of the decide of relative presentation.       Speed:         acrossity, (released at low velocity into zone of active generation)       0.51 m/s         direct spray, spray painting in shallow bootts, gas discharge (active generation into zone of rapid air motion)       1-2.5 m/s (200-500 l/m)         Within each range       Upper end of the range       1.0 Extra time of a super section of the range       1.2.5 m/s (200-500 l/m)         2. Contramining of low toxicity or of inducation of unique over one of the range       1.0 Extra time of the range       1.0 Extra time of the range         2. Contramining of low toxicity or of inducation and the stratection point (in simple access). Therefore the air speed at the extraction point, and inspeed at the extraction point. A subscript and the extraction point, of a simple access, the induce and induce a subscript induced at the extraction point. A subscript and the extraction point, of a simple access, the extraction point, of a simple access, the extraction access, the extraction point, of a simple access, the extraction point, of a simple access, the extraction access, the extraction access, the extraction access, the extraction access, the extra time access and time extra time access and time extra time access and time extr		General exhaust is adequate under normal conditions. If risk of overexposure exists, wear SAA approved respirator. Correct fit is essential to obtain adequate protection. Provide adequate ventilation in warehouse or closed storage areas. Air contaminants generated in the workplace possess varying "escape" velocities which, in turn, determine the "capture velocities" of fresh					
- Type Containing and.       - Development         - serveds, (released at low velocity into zone of active generation)       -0.5 fm is         - development		throughing an required to enectively remove the contaminant.					
Lade tables (released a flow which y min 2 sho in during generation)       U2-1 min         detect sprs, sprix paining in shallow booths, gas detectings (stative generation into zone of rapid air motion)       1-2.5 m/s (200-500 t/mi         Within each range the appropriate value depends on:       Lower and of the range       Upper and of the range         1: Room all currents minimal of favourable to capity.       2. Contaminants of high toxicity       3. Intermittent, low production.         3: Intermittent, low production.       3: High production (haver years)       4. Simple theory years         4: Large hood or large air mass in motion       4: Simal bood-boal control only         Simple theory shows that air velocity fails apidly with distance away from the opening of a simple extraction pier. Velocity generally decrew with the square of distance from the averaciden pier (in simple cases). Therefore than air special motion       1:		Type of Contaminant: Speed:					
Personal protection <ul> <li>Safety glasses with side abunda.</li> <li>Even af acceptor spin abundation of the stange</li> <li>Room ari currents minimal or favourable to capture</li> <li>I) Disturbing commanication of the stange</li> <li>Room ari currents minimal or favourable to capture</li> <li>I) Disturbing commanication of the stange</li> <li>Room ari currents minimal or favourable to capture</li> <li>I) Disturbing commanication of the stange</li> <li>Room ari currents minimal or favourable to capture</li> <li>I) Disturbing commanication of the stange of the</li></ul>		direct spray, spray, painting in shallow booths, gas dischare	ration)	$1_{-2.5} \text{ m/s} (200_{-500} \text{ f/min})$			
Lower end of the range     Upper end of the range       1     Recom air currents minimal or favourable to capture     1: Disturbing room air currents       2     Contaminants of low toxicity or of nuisance value only     2: Contaminants of low toxicity       3     High production, heavy use       4     Large hood or large air mass in motion     4: Small hood-local currents       3     High production, heavy use       4     Large hood or large air mass in motion     4: Small hood-local currents on joint should be adjusted.       Accounting in the square of distance from the contanianting source. The air velocity at the extraction point. Other mechanical considerations, producing performance deficits within the extraction appertus, make it essential that theoretical air velocities are multiplied factors of 10 or more when extraction systems are installed or used.       Personal protection     Image: Status of the strange on tabulant the extraction systems are installed or used.       Eye and face protection <ul> <li>Stafey glasses with side shields.</li> <li>Chemical goggies.</li> <li>Cost titing goggies.</li> <li>Stafey glasses with side shields.</li> <li>Chemical goggies.</li> <li>Cost titing goggies.</li> <li>Stafey glasses with side shields.</li> <li>Chemical equipment should be readial valiable.</li> <li>Cost titing gas glipt goggies</li> <li>Stain protection</li> <li>Stafey glasses, such ashould schemowel at the fit signs of gray erefr</li></ul>		Within each range the appropriate value depends on:		1-2.3 11/3 (200-300 1/11111.)			
1: Room air currents minimal or favourable to capture       1: Disturbing room air currents         1: Contaminants of low toxicity or fusions value only.       2: Contaminants of high toxicity.         1: Large hood or large air mass in motion       4: Small hood-local control only.         Simple theory shows that al viceolity (alls rapid) with distance away from the opening of a simple extraction pipe. Velocity generally deere with the square of distance from the contaminating source. The air velocity at the extraction pipe. Velocity generally deere with the square of distance from the contaminating source. The air velocity at the extraction pipe. Velocity generally deere with the square of distance from the contaminating source. The air velocity at the extraction pipe. Velocity generally deere distance from the contaminating source. The air velocity at the extraction pipe. Velocity generally deere distance from the contaminating source. The air velocity at the extraction pipe. Velocity generally deere distance from the contaminating source. The air velocity at the extraction pipe. Velocity generally deere distance from the contaminating source. The air velocity at the extraction pipe. Velocity generally deere distance from the contaminating source. The air velocity at the extraction pipe. Velocity generally deere distance from the contaminating source. The air velocity at the extraction pipe. Velocity generally deere distance from the contaminating of the straction pipe. Velocity at the extraction pipe. Velocity generally deere distance from the extraction pipe. Velocity at the extraction pipe. Velocity and the optication of the velocity of pipe source.         Personal protection       • Safety glasses with side shields.       • Safety glasses with side shields.         • Contrad integer protection on us		Lower end of the range	Upper end of the range				
<ul> <li>2. Contaminants of low toxicity or of nuisance value only.</li> <li>2. Contaminants of high toxicity         <ul> <li>3. High production.</li> <li>3. High production have value</li> <li>4. Small hood-local control only</li> </ul> </li> <li>Simple theory shows that air velocity falls rapidly with distance away from the opening of a simple extraction pipe. Velocity generally decreating the extraction point. Other mechanical considerations, producing performance deficits within the extraction paper. Velocity and the advance of non-toxicity of or extraction point. Other mechanical considerations, producing performance deficits within the extraction apparatus, make it essential that theoretical air velocities are multiplier factors of 10 or more when extraction systems are installed or used.</li> <li>Personal protection</li> <li>Safety glasses with side shields.</li> <li>Cobemical goggies.</li> <li>Safety glasses with side shields.</li> <li>Cobemical goggies.</li> <li>Contract inters on pose a special hazard; soft contact interses may absorb and concentrate initiants. A written policy document, description of the cases on aspeciate the consolid the groups on task. The Safety glasses with side shields.</li> <li>Cobemical goggies.</li> <li>Safety glasses with side shields.</li> <li>Cobemical goggies.</li> <li>Safety glasses on esticitans on use, should be readive available. In the event of themical exposure, begin eye irrigation immediately irremove contact lines as on a special hazard; soft contact lines as on a special hazard; soft contact lines as should be removed at the first sign of periods.</li> <li>Contact lines as son as paractale. Luces should be removed at the first sign of periods soft microstantely intervents have washed hands thoroughly. [CDC NIOSH Current Intelligence Bulletin 58], (ASINZS 1336 in attraction on the social glassifistin goggies.</li> <li>Start protection</li> <li>Stare</li></ul>		1: Room air currents minimal or favourable to capture	1: Disturbing room air currents				
3: Intermittent, low production.       3: High production, heavy use         4: Large hood or large air mass in motion       4: Small hood-local control only         Simple theory shows that air velocity falls rangibly with distance away from the opening of a simple extraction point. (In simple cases). Therefore the arispaced at the extraction point. Other mechanical constrainting source. The air velocity at the extraction point. Other mechanical constrainting source. The air velocity at the extraction point. Other mechanical constrainting source. The air velocity at the extraction point. Other mechanical constrainting source. The air velocity at the extraction point. Other mechanical constrainting source. The air velocity at the extraction point. Other mechanical constrainting source. The air velocity attent the extraction point. Other mechanical constrainting source. The air velocity attent the extraction point. Other mechanical constraints in source. The air velocities are multiplies factors of 10 or more when extraction systems are installed or used.         Personal protection       Safety glasses with side shields.         Contract lenses may pose a special hazard, soft contract lenses may absorb and concentrate initiants. A written policy document, descript the water and and and a social to readive valuables. In the event of chemical exposure, begive at adsorption for the class of chemical source is an account of injury experime. Meclas and Insta-4d personnel should be randing and adsorption. The trans should in crash as should be readive at a social to runnot and (shifted personnel should be randing and adsorption. The trans about be randing and adsorption. The trans about be randing and adsorption. The trans about an account of injury experime. Meclas and Insta-4d personnel should be randing and a quovalent.         E		2: Contaminants of low toxicity or of nuisance value only.	2: Contaminants of high toxicity				
4: Large hood or large air mass in motion       4: Small hood-local control only         Simple theory shows that air velocity fails rapidly with distance away from the opening of a simple extraction point should be adjusted, accordingly, after reference to distance from the containing source. The air velocity after attencion point should be adjusted, accordingly, after reference to distance from the containing source. The air velocity after reference to distance from the containing source. The air velocity after reference to distance from the containing source. The air velocity after reference to distance from the containing source. The air velocity after reference to distance from the containing source. The air velocity after reference to distance from the containing source. The air velocity after reference to distance from the containing source. The air velocity after reference to distance from the containing source. The air velocity after reference to distance from the containing source. The air velocity after reference to distance from the extraction paperatus, make it essential that theoretical air velocities are multiplies factors of 10 or more when extraction systems are installed or used.         Personal protection       VOVO		3: Intermittent, low production.	3: High production, heavy use				
Simple theory shows that air velocity fails rapidly with distance away from the pering of a simple extraction pipe. Velocity generally decre- with the square of distance from the contaminating source. The air speed at the extraction print. Other mechanical considerations, producing performance deficits within the extraction approximation point. Other mechanical considerations, producing performance deficits within the extraction approximation point. Other mechanical considerations, producing performance deficits within the extraction apparatus, make it essential that theoretical air velocities are multiplied factors of 10 or more when extraction systems are installed or used.         Personal protection       Safety glasses with side shields.         • Chemical goggles.       • Safety glasses with side shields.         • Chemical goggles.       • Chemical goggles.         • Contract lenses may absorb and concentrate initiants. A written policy document, descri the waring of lenses or restrictions on use, should be created for each workplace or task. This should include a review of lens should be remove contact lens as so con a practical bin use and an account of injury experiment. Modical and first-aid personnel should be trained their removal and suitable equipment should be transed. Lens should be termed. Lens should be terms should be terms and an adsorphin of the class of chemicals in use and an account of injury experiments. Modical and first-aid personnel should be remove contact lens as so con as practical blass hould be termed.         Handsifieet protection       See thand protect shin sensitisation in predisposed individuals. Care must be taken, when removing gloves and other protection equipment, to avoid all prospine termed when handing small quantities.         Handsifieet protec		4: Large hood or large air mass in motion	4: Small hood-local control only				
Personal protection       Image: Constraint of the second se		Simple theory shows that air velocity falls rapidly with distance with the square of distance from the extraction point (in simpl accordingly, after reference to distance from the contaminatin 1-2 m/s (200-400 f/min.) for extraction of solvents generated considerations, producing performance deficits within the ext factors of 10 or more when extraction systems are installed of	se away from the opening of a simple extraction pipe le cases). Therefore the air speed at the extraction p ng source. The air velocity at the extraction fan, for e in a tank 2 meters distant from the extraction point. raction apparatus, make it essential that theoretical or used.	<ul> <li>Velocity generally decreases point should be adjusted, example, should be a minimum of Other mechanical air velocities are multiplied by</li> </ul>			
Eye and face protection <ul> <li>Safety glasses with side shields.</li> <li>Chemical goggles.</li> <li>Contact lenses may pose a special hazard; soft contact lenses may absorb and concentrate irritants. A written policy document, description of the class of chemicals in use and an account of injury experience. Medical and first-aid personnel should be trained their removal and suitable equipment should be readily available. In the event of chemical exposure, being new irritation - lens should be trained their removal and suitable equipment should be removed at the first signs of eye redness or irritation - lens should be removed at the first signs of eye redness or irritation - lens should be removed at the first signs of eye redness or irritation - lens should be removed at the first signs of eye redness or irritation - lens should be removed at the first signs of eye redness or irritation - lens should be removed at the first signs of eye redness or irritation - lens should be removed and equivalent]          <ul> <li>Close fitting gas tight goggles</li> <li>Skin protection</li> <li>See Hand protection below</li> </ul> </li> <li>No TEE:         <ul> <li>The material may produce skin sensitisation in predisposed individuals. Care must be taken, when removing gloves and other protection equipment, to avoid all possible skin contact.</li> <li>Contrain tende leather items, such as shoes, belts and watch-bands should be removed and destroyed.</li> <li>No special equipment needed when handling small quantities.</li> <li>OTHERWISE:</li> <li>For potentially moderate exposures:</li> <li>Wear general protective gloves, eg. Itylt weight rubber gloves.</li> <li>For potentially heavy exposures:</li> <li>Wear general protective gloves, eg. PVC. and safety footwear.&lt;</li></ul></li></ul>	Personal protection						
Skin protection       See Hand protection below         NOTE: <ul> <li></li></ul>	Eye and face protection	<ul> <li>Safety glasses with side shields.</li> <li>Chemical goggles.</li> <li>Contact lenses may pose a special hazard; soft contact lenses may absorb and concentrate irritants. A written policy document, describing the wearing of lenses or restrictions on use, should be created for each workplace or task. This should include a review of lens absorption and adsorption for the class of chemicals in use and an account of injury experience. Medical and first-aid personnel should be trained in their removal and suitable equipment should be readily available. In the event of chemical exposure, begin eye irrigation immediately and remove contact lens as soon as practicable. Lens should be removed at the first signs of eye redness or irritation - lens should be removed in a clean environment only after workers have washed hands thoroughly. [CDC NIOSH Current Intelligence Bulletin 59], [AS/NZS 1336 or national equivalent]</li> </ul>					
Hands/feet protection       NOTE:            Hands/feet protection           Herminated leather items, such as shoes, belts and watch-bands should be removed and destroyed.             Hands/feet protection           Contaminated leather items, such as shoes, belts and watch-bands should be removed and destroyed.             Hands/feet protection           Contaminated leather items, such as shoes, belts and watch-bands should be removed and destroyed.             Hands/feet protection           OtherBRWISE:             For potentially moderate exposures:           Year general protective gloves, eg. light weight rubber gloves.             For potentially heavy exposures:           Year demical protective gloves, eg. PVC. and safety footwear.             Body protection           See Other protection below             No special equipment needed when handling small quantities.           Other protection below             Other protection           No special equipment needed when handling small quantities.             Other protection           No special equipment needed when handling small quantities.             Other protection             Do as spray on hot surfaces.             Se oth spray on hot surfaces.             <	Skin protection	See Hand protection below					
Body protection       See Other protection below         No special equipment needed when handling small quantities.       OTHERWISE:         Overalls.       > Skin cleansing cream.         Eyewash unit.       > Do not spray on hot surfaces.         Do not spray on hot surfaces.       > The clothing worn by process operators insulated from earth may develop static charges for higher (up to 100 times) than the minimum	Hands/feet protection	<ul> <li>NOTE:</li> <li>The material may produce skin sensitisation in predisposed individuals. Care must be taken, when removing gloves and other protective equipment, to avoid all possible skin contact.</li> <li>Contaminated leather items, such as shoes, belts and watch-bands should be removed and destroyed.</li> <li>No special equipment needed when handling small quantities.</li> <li>OTHERWISE:</li> <li>For potentially moderate exposures:</li> <li>Wear general protective gloves, eg. light weight rubber gloves.</li> <li>For potentially heavy exposures:</li> <li>Wear chemical protective gloves, eg. PVC. and safety footwear.</li> </ul>					
Other protection       No special equipment needed when handling small quantities.         Other protection       • Overalls.         • Eyewash unit.       • Eyewash unit.         • Do not spray on hot surfaces.       • The clothing worn by process operators insulated from earth may develop static charges far higher (up to 100 times) than the minimum	Body protection	See Other protection below					
ignition energies for various flammable gas-air mixtures. This holds true for a wide range of clothing materials including cotton. Avoid dangerous levels of charge by ensuring a low resistivity of the surface material worn outermost.	Other protection	No special equipment needed when handling small quantities. OTHERWISE: • Overalls. • Skin cleansing cream. • Eyewash unit. • Do not spray on hot surfaces. • The clothing worn by process operators insulated from earth may develop static charges far higher (up to 100 times) than the minimum ignition energies for various flammable gas-air mixtures. This holds true for a wide range of clothing materials including cotton. • Avoid dangerous levels of charge by ensuring a low resistivity of the surface material worn outermost.					

# **Respiratory protection**

Type AX-P Filter of sufficient capacity. (AS/NZS 1716 & 1715, EN 143:2000 & 149:2001, ANSI Z88 or national equivalent)

Where the concentration of gas/particulates in the breathing zone, approaches or exceeds the "Exposure Standard" (or ES), respiratory protection is required. Degree of protection varies with both face-piece and Class of filter; the nature of protection varies with Type of filter.

Required Minimum Protection Factor	Half-Face Respirator	Full-Face Respirator	Powered Air Respirator
up to 10 x ES	AX-AUS P2	-	AX-PAPR-AUS / Class 1 P2
up to 50 x ES	-	AX-AUS / Class 1 P2	-
up to 100 x ES	-	AX-2 P2	AX-PAPR-2 P2 ^

A(All classes) = Organic vapours, B AUS or B1 = Acid gasses, B2 = Acid gas or hydrogen cyanide(HCN), B3 = Acid gas or hydrogen cyanide(HCN), E = Sulfur dioxide(SO2), G = Agricultural chemicals, K = Ammonia(NH3), Hg = Mercury, NO = Oxides of nitrogen, MB = Methyl bromide, AX = Low boiling point organic compounds(below 65 degC)

- Cartridge respirators should never be used for emergency ingress or in areas of unknown vapour concentrations or oxygen content.
- The wearer must be warned to leave the contaminated area immediately on detecting any odours through the respirator. The odour may indicate that the mask is not functioning properly, that the vapour concentration is too high, or that the mask is not properly fitted. Because of these limitations, only restricted use of cartridge respirators is considered appropriate.
- Cartridge performance is affected by humidity. Cartridges should be changed after 2 hr of continuous use unless it is determined that the humidity is less than 75%, in which case, cartridges can be used for 4 hr. Used cartridges should be discarded daily, regardless of the length of time used

Generally not applicable.

Aerosols, in common with most vapours/ mists, should never be used in confined spaces without adequate ventilation. Aerosols, containing agents designed to enhance or mask smell, have triggered allergic reactions in predisposed individuals.

## **SECTION 9** Physical and chemical properties

#### Information on basic physical and chemical properties

Appearance	Light golden yellow flammable liquid with a solvent odour; not miscible with water. Supplied as an aerosol pack. Contents under <b>PRESSURE</b> . Contains highly flammable hydrocarbon propellant.			
Physical state	Liquid	Relative density (Water = 1)	0.65-0.73	
Odour	Not Available	Partition coefficient n-octanol / water	Not Available	
Odour threshold	Not Available	Auto-ignition temperature (°C)	Not Available	
pH (as supplied)	Not Applicable	Decomposition temperature	Not Available	
Melting point / freezing point (°C)	Not Available	Viscosity (cSt)	Not Available	
Initial boiling point and boiling range (°C)	Not Available	Molecular weight (g/mol)	Not Applicable	
Flash point (°C)	Not Available	Taste	Not Available	
Evaporation rate	Not Available	Explosive properties	Not Available	
Flammability	Not Available	Oxidising properties	Not Available	
Upper Explosive Limit (%)	Not Available	Surface Tension (dyn/cm or mN/m)	Not Available	
Lower Explosive Limit (%)	Not Available	Volatile Component (%vol)	>60	
Vapour pressure (kPa)	Not Available	Gas group	Not Available	
Solubility in water	Immiscible	pH as a solution (Not Available%)	Not Applicable	
Vapour density (Air = 1)	Not Available	VOC g/L	Not Available	

#### **SECTION 10 Stability and reactivity**

Reactivity	See section 7
Chemical stability	<ul> <li>Elevated temperatures.</li> <li>Presence of open flame.</li> <li>Product is considered stable.</li> <li>Hazardous polymerisation will not occur.</li> <li>Presence of heat source</li> <li>Presence of an ignition source</li> </ul>
Possibility of hazardous reactions	See section 7
Conditions to avoid	See section 7
Incompatible materials	See section 7
Hazardous decomposition products	See section 5

## **SECTION 11 Toxicological information**

## Information on toxicological effects

	Inhalation of vapours may cause drowsiness and dizziness. This may be accompanied by sleepiness, reduced alertness, loss of reflexes, lack of co-ordination, and vertigo.
	Inhalation of aerosols (mists, fumes), generated by the material during the course of normal handling, may be damaging to the health of the individual.
	There is some evidence to suggest that the material can cause respiratory irritation in some persons. The body's response to such irritation can cause further lung damage.
	Inhalation of toxic gases may cause:
	Central Nervous System effects including depression, headache, confusion, dizziness, stupor, coma and seizures;
Inhaled	respiratory: acute lung swellings, shortness of breath, wheezing, rapid breathing, other symptoms and respiratory arrest;
	heart: collapse, irregular heartbeats and cardiac arrest;
	gastrointestinal: irritation, ulcers, nausea and vomiting (may be bloody), and abdominal pain.
	Inhalation of high concentrations of gas/vapour causes lung irritation with coughing and nausea, central nervous depression with headache and dizziness, slowing of reflexes, fatigue and inco-ordination.
	Central nervous system (CNS) depression may include general discomfort, symptoms of giddiness, headache, dizziness, nausea, anaesthetic effects, slowed reaction time, slurred speech and may progress to unconsciousness. Serious poisonings may result in respiratory depression and may be fatal.

	Material is highly volatile and may quickly form a concentrated atmosphere in confined or unventilated areas. The vapour may displace and replace air in breathing zone, acting as a simple asphyxiant. This may happen with little warning of overexposure. WARNING:Intentional misuse by concentrating/inhaling contents may be lethal.			
Ingestion	Accidental ingestion of the material may be damaging to the health of the individual. Not normally a hazard due to physical form of product. Considered an unlikely route of entry in commercial/industrial environments Central nervous system (CNS) depression may include general discomfort, symptoms of giddiness, headache, dizziness, nausea, anaesthetic effects, slowed reaction time, slurred speech and may progress to unconsciousness. Serious poisonings may result in respiratory depression and may be fatal.			
Skin Contact	Repeated exposure may cause skin cracking, flaking or drying following Spray mist may produce discomfort Open cuts, abraded or irritated skin should not be exposed to this materia The material may accentuate any pre-existing dermatitis condition	normal handling and use. al		
Eye	This material can cause eye irritation and damage in some persons. Not	considered to be a risk because of the extreme volatility of the gas.		
Chronic	Substance accumulation, in the human body, may occur and may cause some concern following repeated or long-term occupational exposure. There is some evidence that inhaling this product is more likely to cause a sensitisation reaction in some persons compared to the general population. There is limited evidence that, skin contact with this product is more likely to cause a sensitisation reaction in some persons compared to the general population. Main route of exposure to the gas in the workplace is by inhalation. Oil may contact the skin or be inhaled. Extended exposure can lead to eczema, inflammation of hair follicles, pigmentation of the face and warts on the soles of the feet. Constant or exposure over long periods to mixed hydrocarbons may produce stupor with dizziness, weakness and visual disturbance, weight loss and anaemia, and reduced liver and kidney function. Skin exposure may result in drying and cracking and redness of the skin.			
Dy-Mark Protech Chain	ΤΟΧΙΟΙΤΥ	IRRITATION		
Lubricant	Not Available	Not Available		
	ΤΟΧΙΟΙΤΥ	IRRITATION		
white mineral oil (petroleum)	Dermal (rabbit) LD50: >2000 mg/kg <sup>[1]</sup>	Eye: no adverse effect observed (not irritating) <sup>[1]</sup>		
	Inhalation(Rat) LC50; >4.5 mg/l4h <sup>[1]</sup>	Skin: adverse effect observed (irritating) <sup>[1]</sup>		
	Oral (Rat) LD50; >5000 mg/kg <sup>[1]</sup>	Skin: no adverse effect observed (not irritating) <sup>[1]</sup>		
2-methylpentane	ΤΟΧΙΟΙΤΥ	IRRITATION		
	Oral (Rat) LD50; ~15.84 mg/kg <sup>[1]</sup>	Not Available		
LPG (liquefied petroleum gas)	ΤΟΧΙΟΙΤΥ	IRRITATION		
	Inhalation(Rat) LC50; 658 mg/l4hl <sup>2]</sup>	Not Available		
Legend:	1. Value obtained from Europe ECHA Registered Substances - Acute toxicity 2.* Value obtained from manufacturer's SDS. Unless otherwise specified data extracted from RTECS - Register of Toxic Effect of chemical Substances			
Dy-Mark Protech Chain Lubricant	Animal studies indicate that normal, branched and cyclic paraffins are absorbed from the gastrointestinal tract and that the absorption of n-paraffins is inversely proportional to the carbon chain length, with little absorption above C30. With respect to the carbon chain lengths likely to be present in mineral oil, n-paraffins may be absorbed to a greater extent than iso- or cyclo-paraffins. The major classes of hydrocarbons are well absorbed into the gastrointestinal tract in various species. In many cases, the hydrophobic hydrocarbons are ingested in association with fats in the diet. Some hydrocarbons may appear unchanged as in the lipoprotein particles in the gut lymph, but most hydrocarbons partly separate from fats and undergo metabolism in the gut cell. The gut cell may play a major role in determining the proportion of hydrocarbon that becomes available to be deposited unchanged in peripheral tissues such as in the body fat stores or the liver.			
WHITE MINERAL OIL (PETROLEUM)	Oral (rat) TCLo: 92000 mg/kg/92D-Cont. Generally the toxicity and irritation is of low order. White oils and highly/solvent refined oils have not shown the long term risk of skin cancer that follows persistent skin contamination with some other mineral oils, due in all probability to refining that produces low content of both polyaromatics (PAH) and benz-alpha-pyrenes (BaP) The substance is classified by IARC as Group 3: <b>NOT</b> classifiable as to its carcinogenicity to humans. Evidence of carcinogenicity may be inadequate or limited in animal testing.			
Dy-Mark Protech Chain Lubricant & 2-METHYLPENTANE & LPG (LIQUEFIED PETROLEUM GAS)	No significant acute toxicological data identified in literature search.			
Dy-Mark Protech Chain Lubricant & LPG (LIQUEFIED PETROLEUM GAS)	inhalation of the gas			
Dy-Mark Protech Chain Lubricant & WHITE MINERAL OIL (PETROLEUM)	The materials included in the Lubricating Base Oils category are related from both process and physical-chemical perspectives; The potential toxicity of a specific distillate base oil is inversely related to the severity or extent of processing the oil has undergone, since: • The adverse effects of these materials are associated with undesirable components, and • The levels of the undesirable components are inversely related to the degree of processing; • Distillate base oils receiving the same degree or extent of processing will have similar toxicities; • The potential toxicity of residual base oils is independent of the degree of processing the oil receives. • The reproductive and developmental toxicity of the distillate base oils is inversely related to the degree of processing. Unrefined & mildly refined distillate base oils contain the highest levels of undesirable components, have the largest variation of hydrocarbon molecules and have shown the highest potential cancer-causing and mutation-causing activities. Highly and severely refined distillate base oils by removing or transforming undesirable components. In comparison to unrefined and mildly refined base oils, the highly and severely refined distillate base oils have a smaller range of hydrocarbon molecules and have demonstrated very low mammalian toxicity. Testing of residual oils for mutation-causing and cancer-causing potential has shown negative results, supporting the belief that these materials lack biologically active components or the components are largely non-bioavailable due to their molecular size.			

Continued...

Toxicity testing has consistently shown that lubricating base oils have low acute toxicities. Numerous tests have shown that a lubricating base oil s mutagenic and carcinogenic potential correlates with its 3-7 ring polycyclic aromatic compound (PAC) content, and the level of DMSO extractables (e.g. IP346 assay), both characteristics that are directly related to the degree/conditions of processing.

For highly and severely refined distillate base oils:

In animal studies, the acute, oral, semilethal dose is >5g/kg body weight and the semilethal dose by skin contact is >2g/kg body weight. The semilethal concentration for inhalation is 2.18 to >4 mg/L. The materials have varied from "non-irritating" to "moderately irritating" when tested for skin and eye irritation. Testing for sensitisation has been negative. The effects of repeated exposure vary by species; in animals, effects to the testes and lung have been observed, as well as the formation of granulomas. In animals, these substances have not been found to cause reproductive toxicity or significant increases in birth defects. They are also not considered to cause cancer, mutations or chromosome aberrations.

Acute Toxicity	×	Carcinogenicity	×
Skin Irritation/Corrosion	×	Reproductivity	×
Serious Eye Damage/Irritation	×	STOT - Single Exposure	×
Respiratory or Skin sensitisation	×	STOT - Repeated Exposure	×
Mutagenicity	×	Aspiration Hazard	×
		Legend: 🗙 – Data either r	ot available or does not fill the criteria for classification

X – Data either not

Data available to make classification

#### **SECTION 12 Ecological information**

Toxicity

#### Endpoint Test Duration (hr) Species Value Source **Dy-Mark Protech Chain** Not Not Not Lubricant Not Available Not Available Available Available Available Test Duration (hr) Species Value Source Endpoint white mineral oil (petroleum) LC50 96h Fish >10000mg/L 2 Endpoint Test Duration (hr) Species Value Source 2-methylpentane EC50(ECx) 96h Algae or other aquatic plants 4.321mg/l 2 EC50 96h Algae or other aquatic plants 4.321mg/l 2 Endpoint Test Duration (hr) Species Value Source EC50(ECx) 96h Algae or other aquatic plants 7.71mg/l 2 LPG (liquefied petroleum gas) LC50 24.11mg/l 96h Fish 2 EC50 96h Algae or other aquatic plants 7.71ma/l 2 Extracted from 1. IUCLID Toxicity Data 2. Europe ECHA Registered Substances - Ecotoxicological Information - Aquatic Toxicity 4. US EPA, Legend: Ecotox database - Aquatic Toxicity Data 5. ECETOC Aquatic Hazard Assessment Data 6. NITE (Japan) - Bioconcentration Data 7. METI (Japan) - Bioconcentration Data 8. Vendor Data

For Petroleum Hydrocarbon Gases

Environmental Fate: Petroleum hydrocarbon gases are primarily produced in petroleum refineries, or in gas plants that separate natural gas and natural gas liquids. This category contains 99 petroleum hydrocarbon gas substances, the majority of which never reach the consumer. Petroleum hydrocarbon gases do not contain inorganic compounds, (e.g. hydrogen sulfide, ammonia, and carbon monoxide), other than asphyxiant gases; the low molecular weight hydrocarbon molecules are primarily responsible for the hazard associated with these gases.

Atmospheric Fate: All components of these gases will evaporate to the air where interaction with hydroxyl radicals is an important fate process. Substances in refinery gases that evaporate to air may undergo indirect, gas-phase oxidation reaction with hydroxyl radicals and this is an important fate process for these substances. Half-lives for refinery gases range from 960 days, (methane), to 0.16 days, (butadiene). The constituents of the C5- C6 hydrocarbon gases have light breakdown half-lives of approximately two days. The inorganic gases are chemically stable and may be lost to the atmosphere or simply become involved in the environmental recycling of their atoms.

Terrestrial Fate: Biological breakdown of these organisms is not expected to be an important fate process since they tend to evaporate to the air, however, some of the higher weight components may become available for microbial attack. Naphtha gases are also considered to be inherently biodegradable.

Aquatic Fate: The solubilities of these substances in water vary, ranging from approximately 22 parts per million to several hundred parts per million. Some of these gasses have substantial water solubility, but they will eventually evaporate to the atmosphere. Refinery gases are not broken down by water but, they will be broken down by microbes. Gaseous hydrocarbons are widespread in nature and numerous types of microbes have evolved which are capable of oxidizing these substances as their sole energy source. Ecotoxicity: These substances vary in their toxicities to aquatic organisms from slightly toxic to moderately toxic. They are not expected to persist long enough in the environment to elicit toxicity. Emissions of petroleum hydrocarbon gases to the atmosphere would not likely result in acutely toxic concentrations in adjacent water bodies because such emissions will tend to remain in the atmosphere. Several of the constituents in refinery gases were shown to be highly hazardous to aquatic organisms in laboratory toxicity tests where exposure concentrations can be maintained over time. Hydrogen sulfide was shown to be the most toxic constituent to fish, and invertebrates.

## for lubricating oil base stocks:

Vapor Pressure Vapor pressures of lubricating base oils are reported to be negligible. In one study, the experimentally measured vapour pressure of a solvent-dewaxed heavy paraffinic distillate base oil was 1.7 x 10exp-4 Pa . Since base oils are mixtures of C15 to C50 paraffinic, naphthenic, and aromatic hydrocarbon isomers, representative components of those structures were selected to calculate a range of vapor pressure. The estimated vapor pressure values for these selected components of base oils ranged from 4.5 x 10exp-1 Pa to 2 x 10exp-13Pa. Based on Dalton's Law the expected total vapour pressure for base oils would fall well below minimum levels (10exp-5 Pa) of recommended experimental procedures.

Partition Coefficient (log Kow): In mixtures such as the base oils, the percent distribution of the hydrocarbon groups (i.e., paraffins, naphthenes, and aromatics) and the carbon chain lengths determines in-part the partitioning characteristics of the mixture. Generally, hydrocarbon chains with fewer carbon atoms tend to have lower partition coefficients than those with higher carbon numbers. However, due to their complex composition, unequivocal determination of the log Kow of these hydrocarbon mixtures cannot be made. Rather, partition coefficients of selected C15 chain-length hydrocarbon structures representing paraffinic, naphthenic, and aromatic constituents in base oil lubricants were modelled. Results showed typical log Kow values from 4.9 to 7.7, which were consistent with values of >4 for lubricating oil basestocks

Water Solubility:When released to water, base oils will float and spread at a rate that is viscosity dependent. While water solubility of base oils is typically very low, individual hydrocarbons exhibit a wide range of solubility depending on molecular weight and degree of unsaturation. Decreasing molecular weight (i.e., carbon number) and increasing levels of unsaturation increases the water solubility of these materials. As noted for partition coefficient, the water solubility of lubricating base oils cannot be determined due to their complex mixture characteristics. Therefore, the water solubility of individual C15 hydrocarbons representing the different groups making up base oils (i.e., linear and branched paraffins, naphthenes, and aromatics) was modelled. Based on water solubility modelling of those groups, aqueous solubilities are typically much less than 1 ppm. (0.003-0.63 mg/l)

#### **Environmental Fate:**

Photodegradation: Chemicals having potential to photolyse have UV/visible absorption maxima in the range of 290 to 800 nm. Some chemicals have absorption maxima significantly below 290 nm and consequently cannot undergo direct photolysis in sunlight (e.g. chemicals such as alkanes, alkenes, alkenes, saturated alcohols, and saturated acids). Most hydrocarbon constituents of the materials in this category are not expected to photolyse since they do not show absorbance within the 290-800 nm range. However, photodegradation of polyaromatic hydrocarbons (PAHs) can occur and may be a significant degradation pathway for these constituents of lubricating base oils. The degree and rate at which PAHs may photodegrade depend upon whether conditions allow penetration of light with sufficient energy to effect a change. For example, polycyclic aromatic compounds (PAC) compounds bound to sediments may persist due to a lack of sufficient light penetration

Atmospheric gas-phase reactions can occur between organic chemicals and reactive molecules such as photochemically produced hydroxyl radicals, ozone and nitrogen oxides. Atmospheric oxidation as a result of radical attack is not direct photochemical degradation, but indirect degradation. In general, lubricating base oils have low vapour pressures and volatilisation is not expected to be a significant removal mechanism for the majority of the hydrocarbon components. However, some components (e.g., C15 branched paraffins and naphthenes) appear to have the potential to volatilise Atmospheric half-lives of 0.10 to 0.66 days have been calculated for representative C15 hydrocarbon components of lubricating base oils

Stability in Water: Chemicals that have a potential to hydrolyze include alkyl halides, amides, carbamates, carboxylic acid esters and lactones, epoxides, phosphate esters, and sulfonic acid esters. Because lubricating base oils do not contain significant levels of these functional groups, materials in the lubricating base oils category are not subject to hydrolysis

Chemical Transport and Distribution in the Environment : Based on the physical-chemical characteristics of component hydrocarbons in lubricating base oils, the lower molecular weight components are expected to have the highest vapour pressures and water solubilities, and the lowest partition coefficients. These factors enhance the potential for widespread distribution in the environment. To gain an understanding of the potential transport and distribution of lubricating base oil components, the EQC (Equilibrium Criterion) model was used to characterize the environmental distribution of different C15 compounds representing different structures found in lube oils (e.g., paraffins, naphthenes, and aromatics). The modelling found partitioning to soil or air is the ultimate fate of these C15 compounds. Aromatic compounds partition principally to soil. Linear paraffins partition mostly to soil, while branching appears to allow greater distribution to air. Naphthenes distribute to both soil and air, with increasing proportions in soil for components with the greater number of ring structures. Because the modelling does not take into account degradation factors, levels modelled in the atmosphere are likely overstated in light of the tendency for indirect photodegradation to occur.

**Biodegradation:** The extent of biodegradation measured for a particular lubricating oil basestock is dependent not only on the procedure used but also on how the sample is presented in the biodegradation test. Lubricant base oils typically are not readily biodegradable in standard 28-day tests. However, since the oils consist primarily of hydrocarbons that are ultimately assimilated by microorganisms, and therefore inherently biodegradable. Twenty-eight biodegradability studies have been reported for a variety of lubricating base oils. Based on the results of ultimate biodegradability tests using modified Sturm and manometric respirometry testing the base oils are expected to be, for the most part, inherently biodegradable. Biodegradable. Biodegradable. Biodegradable. Biodegradable in standard 28-day tests from the manometric respirometry tests on similar materials showed biodegradable. Biodegradation rates from 31 to 50%. Biodegradation rates measured in 21-day CEC tests for similar materials ranged from 13 to 79%.

Numerous acute studies covering fish, invertebrates, and algae have been conducted to assess the ecotoxicity of various lubricating base oils. None of these studies have shown evidence of acute toxicity to aquatic organisms. Eight, 7-day exposure studies using rainbow trout failed to demonstrate toxicity when tested up to the maximum concentration of 1000 mg/L applied as dispersions. Three, 96-hour tests with rainbow trout also failed to show any toxic effects when tested up to 1000 mg/L applied as dispersions. Similarly, three 96-hour tests with rainbow trout also failed to show any toxic effects when tested up to 1000 mg/L applied as dispersions. Similarly, three 96-hour tests with fathead minnows at a maximum test concentration of 1000 mg/L water accommodated fractions (WAF) showed no adverse effects. Two species of aquatic invertebrates (Daphnia magna and Gammarus sp.) were exposed to WAF solutions up to 10,000 mg/L for 48 and 96-hours, respectively, with no adverse effects being observed. Four-day exposures of the freshwater green alga (Scenedesmus subspicatus) to 500 mg/L WAF solutions failed to show adverse effects on growth rate and algal cell densities in four studies Multiple chronic ecotoxicity studies have shown no adverse effects to daphnid survival or reproduction. In 10 of 11 chronic studies, daphnids were exposed for 21 days to WAF preparations of lubricating base oils with no ill effects on survival or reproduction at the maximum concentration of 1000 mg/L. One test detected a reduction in reproduction at 1000 mg/L. Additional data support findings of no chronic toxicity to aquatic invertebrates and fish. No observed effect levels ranged from 550 to 5,000 mg/L when tested as either dispersions or WAFs.

The data described above are supported by studies on a homologous series of alkanes. The author concluded that the water solubility of carbon chains .C10 is too limited to elicit acute toxicity. This also was shown for alkylbenzene compounds having carbon numbers .C15. Since base oils consist of carbon compounds of C15 to C50, component hydrocarbons that are of acute toxicological concern are, for the most part, absent in these materials. Similarly, due to their low solubility, the alkylated two to three ring polyaromatic components in base oils are not expected to cause acute or chronic toxicity. This lack of toxicity is borne out in the results of the reported studies. The effects of crude and refined oils on organisms found in fresh and sea water ha been extensively reviewed.

sea water. Where spillages occur the non-mobile species suffer the greatest mortality, whereas fish species can often escape from the affected region. The extent of the initial mortality depends on the chemical nature of the oil, the location, and the physical conditions, particularly the temperature and wind velocity. Most affected freshwater and marine communities recover from the effects of an oil spill within a year. The occurrence of biogenic hydrocarbons in the world's oceans is well recorded. They have the characteristic isoprenoid structure, and measurements made in water columns indicate a background concentration of 1.0 to 10 ul/l. The higher molecular weight materials are dispersed as particles, with the highest concentrations of about 20 ul/l occurring in the top 3 mm layer of water.

A wide variation in the response of organisms to oil exposures has been noted. The larvae of fish and crustaceans appear to be most susceptible to the water-soluble fraction of crude oil. Exposures of plankton and algae have indicated that certain species of diatoms and green algae are inhibited, whereas microflagellates are not.

For the most part, molluscs and most intertidal worm species appear to be tolerant of oil contamination.

For n-Hexane: Log Kow: 3.17-3.94; Henry s Law Constant: 1.69 atm-m3 mol; Vapor Pressure: 150 mm Hg @ 25 C; Log Koc: 2.90 to 3.61. BOD 5, (if unstated): 2.21; COD: 0.04; ThOD: 3.52.

Atmospheric Fate: n-Hexane is not expected to be directly broken down by sunlight. The main atmospheric removal mechanism is through reactions with hydroxyl radicals, with an approximant half-life of 2.9 days. The smog-producing potential of n-hexane is very low, compared to other alkanes, or chlorinated VOCs. Hydroxyl ion reactions in the upper troposphere, therefore, are probably the primary mechanisms for n-hexane degradation in the atmosphere.

Terrestrial Fate: Surface evaporation is expected to be the main fate process of this substance in soil. The substance has a moderate ability to sorb to soil particles but, is expected to have low potential for leaching into the lower soil depths. n-Hexane is expected to generally stay near the soil surface and, if not appreciably sorbed into the soil matrix, will eventually evaporate. Exceptions would involve locations with shallow groundwater tables where large spills occur - in such cases, n-hexane would spread out to contaminate a large volume of soil. Once introduced into groundwater, n-hexane may be fairly persistent, since its degradation by water is slow and opportunities for biodegradation may be limited, (due to low oxygen conditions), or, where nutrients, such as nitrogen or phosphorus, are in limited supply. Biological breakdown is probably the most significant degradation mechanism in groundwater. Pseudomonas mendocina bacteria have been shown to break the substance down in groundwater and mixed/pure bacterial cultures can utilize the substance, in the presence of oxygen. The most important biological breakdown process involves the conversion of n-hexane to primary alcohols, aldehydes and, ultimately, into fatty acids. In general, unless the n-hexane is buried at some depth within a soil or sediment, evaporation is generally assumed to occur at a much more rapid rate than chemical or biochemical degradation processes.

Aquatic Fate: The dominant transport process from water is evaporation, with an estimated half-life of <3 hours. For standing bodies of water, a half-life no longer than 6.8 days is estimated. The substance has very low water solubility and is resistant to breakdown by water. Few data exist for the biological breakdown of n-hexane in water, however; this process is not considered to be as rapid as evaporation. N-Hexane may be persistent if released to deep sediment.

Ecotoxicity: This substance is not expected to concentrate/accumulate in aquatic organisms or the food chain. These substances are considered to be the most readily biodegradable fractions in petroleum, particularly when oxygen is present in solution. The substance is moderately toxic to rainbow trout, fathead minnow, bluegill, and Daphnia water fleas.

#### For Propane: Koc 460. log

#### Kow 2.36.

Henry's Law constant of 7.07x10-1 atm-cu m/mole, derived from its vapour pressure, 7150 mm Hg, and water solubility, 62.4 mg/L. Estimated BCF: 13.1. Terrestrial Fate: Propane is expected to have moderate mobility in soil. Volatilization from moist soil surfaces is expected to be an important fate process. Volatilization from dry soil

surfaces is based vapor pressure. Biodegradation may be an important fate process in soil and sediment. Aquatic Fate: Propane is expected to adsorb to suspended solids and sediment. Volatilization from water surfaces is expected and half-lives for a model river and model lake are

estimated to be 41 minutes and 2.6 days, respectively. Biodegradation may not be an important fate process in water. Ecotoxicity: The potential for bioconcentration in aquatic organisms is low.

Atmospheric Fate: Propane is expected to exist solely as a gas in the ambient atmosphere. Gas-phase propane is degraded in the atmosphere by reaction with photochemicallyproduced hydroxyl radicals; the half-life for this reaction in air is estimated to be 14 days and is not expected to be susceptible to direct photolysis by sunlight. **DO NOT** discharge into sewer or waterways.

#### Persistence and degradability

Ingredient

Persistence: Water/Soil

Persistence: Air

Continued...

# **Dy-Mark Protech Chain Lubricant**

Ingredient	Persistence: Water/Soil Persistence: Air		
2-methylpentane	LOW	LOW	
Bioaccumulative potential			
Ingredient	Bioaccumulation		
2-methylpentane	LOW (LogKOW = 3.2145)		
Mobility in soil			
Ingredient	Mobility		
2-methylpentane	LOW (KOC = 124.9)		

# **SECTION 13 Disposal considerations**

Waste treatment methods				
Product / Packaging disposal	<ul> <li>DO NOT allow wash water from cleaning or process equipment to enter drains.</li> <li>It may be necessary to collect all wash water for treatment before disposal.</li> <li>In all cases disposal to sewer may be subject to local laws and regulations and these should be considered first.</li> <li>Where in doubt contact the responsible authority.</li> <li>Consult State Land Waste Management Authority for disposal.</li> <li>Discharge contents of damaged aerosol cans at an approved site.</li> <li>Allow small quantities to evaporate.</li> <li>DO NOT incinerate or puncture aerosol cans.</li> <li>Bury residues and emptied aerosol cans at an approved site.</li> </ul>			

# **SECTION 14 Transport information**

Labels Required	
Marine Pollutant	
HAZCHEM	Not Applicable

# Land transport (ADG)

UN number	1950			
UN proper shipping name	AEROSOLS	AEROSOLS		
Transport hazard class(es)	Class2.1SubriskNot Applicable			
Packing group	Not Applicable			
Environmental hazard	Environmentally hazardous			
Special precautions for user	Special provisions	63 190 277 327 344 381 1000ml		

# Air transport (ICAO-IATA / DGR)

UN number	1950			
UN proper shipping name	Aerosols, flammable	Aerosols, flammable		
Transport hazard class(es)	ICAO/IATA Class2.1ICAO / IATA SubriskNot ApplicableERG Code10L			
Packing group	Not Applicable			
Environmental hazard	Environmentally hazardous			
Special precautions for user	Special provisions Cargo Only Packing Instructions Cargo Only Maximum Qty / Pack Passenger and Cargo Packing Instructions Passenger and Cargo Maximum Qty / Pack		A145 A167 A802 203 150 kg 203 75 kg	

Passenger and Cargo Limited Quantity Packing Instructions	Y203
Passenger and Cargo Limited Maximum Qty / Pack	30 kg G

## Sea transport (IMDG-Code / GGVSee)

UN number	1950		
UN proper shipping name	AEROSOLS		
Transport hazard class(es)	IMDG Class     2.1       IMDG Subrisk     Not Applicable		
Packing group	Not Applicable		
Environmental hazard	Marine Pollutant		
Special precautions for user	EMS Number Special provisions Limited Quantities	F-D, S-U 63 190 277 327 344 381 959 1000 ml	

# Transport in bulk according to Annex II of MARPOL and the IBC code

## Not Applicable

Transport in bulk in accordance with MARPOL Annex V and the IMSBC Code

Product name	Group
white mineral oil (petroleum)	Not Available
2-methylpentane	Not Available
LPG (liquefied petroleum gas)	Not Available

# Transport in bulk in accordance with the ICG Code

Product name	Ship Type
white mineral oil (petroleum)	Not Available
2-methylpentane	Not Available
LPG (liquefied petroleum gas)	Not Available

# **SECTION 15 Regulatory information**

#### Safety, health and environmental regulations / legislation specific for the substance or mixture

## white mineral oil (petroleum) is found on the following regulatory lists

Australian Inventory of Industrial Chemicals (AIIC)	International Agency for Research on Cancer (IARC) - Agents Classified by the IARC Monographs
	International Agency for Research on Cancer (IARC) - Agents Classified by the IARC Monographs - Group 1: Carcinogenic to humans
2-methylpentane is found on the following regulatory lists	
Australia Hazardous Chemical Information System (HCIS) - Hazardous Chemicals	Australian Inventory of Industrial Chemicals (AIIC)
LPG (liquefied petroleum gas) is found on the following regulatory lists	
Australia Hazardous Chemical Information System (HCIS) - Hazardous Chemicals	Chemical Footprint Project - Chemicals of High Concern List
Australian Inventory of Industrial Chemicals (AIIC)	

## **National Inventory Status**

National Inventory	Status	
Australia - AIIC / Australia Non-Industrial Use	Yes	
Canada - DSL	Yes	
Canada - NDSL	No (white mineral oil (petroleum); 2-methylpentane; LPG (liquefied petroleum gas))	
China - IECSC	Yes	
Europe - EINEC / ELINCS / NLP	Yes	
Japan - ENCS	No (white mineral oil (petroleum))	
Korea - KECI	Yes	
New Zealand - NZIoC	Yes	
Philippines - PICCS	Yes	
USA - TSCA	Yes	
Taiwan - TCSI	Yes	
Mexico - INSQ	Yes	
Vietnam - NCI	Yes	
Russia - FBEPH	Yes	

National Inventory	Status
Legend:	Yes = All CAS declared ingredients are on the inventory No = One or more of the CAS listed ingredients are not on the inventory. These ingredients may be exempt or will require registration.

## **SECTION 16 Other information**

Revision Date	20/04/2022	
Initial Date	15/09/2014	
SDS Version Summary		
Manatan	Deter of the late	Our Serve He late 1

Version	Date of Update	Sections Updated
10.1	10/12/2021	Classification change due to full database hazard calculation/update.
11.1	20/04/2022	Classification, Ingredients

#### Other information

Classification of the preparation and its individual components has drawn on official and authoritative sources as well as independent review by the Chemwatch Classification committee using available literature references.

The SDS is a Hazard Communication tool and should be used to assist in the Risk Assessment. Many factors determine whether the reported Hazards are Risks in the workplace or other settings. Risks may be determined by reference to Exposures Scenarios. Scale of use, frequency of use and current or available engineering controls must be considered.

#### **Definitions and abbreviations**

PC-TWA: Permissible Concentration-Time Weighted Average PC-STEL: Permissible Concentration-Short Term Exposure Limit IARC: International Agency for Research on Cancer ACGIH: American Conference of Governmental Industrial Hygienists STEL: Short Term Exposure Limit TEEL: Temporary Emergency Exposure Limit。 IDLH: Immediately Dangerous to Life or Health Concentrations ES: Exposure Standard OSF: Odour Safety Factor NOAEL :No Observed Adverse Effect Level LOAEL: Lowest Observed Adverse Effect Level TLV: Threshold Limit Value LOD: Limit Of Detection OTV: Odour Threshold Value BCF: BioConcentration Factors **BEI: Biological Exposure Index** AIIC: Australian Inventory of Industrial Chemicals DSL: Domestic Substances List NDSL: Non-Domestic Substances List IECSC: Inventory of Existing Chemical Substance in China EINECS: European INventory of Existing Commercial chemical Substances ELINCS: European List of Notified Chemical Substances NLP: No-Longer Polymers ENCS: Existing and New Chemical Substances Inventory KECI: Korea Existing Chemicals Inventory NZIoC: New Zealand Inventory of Chemicals PICCS: Philippine Inventory of Chemicals and Chemical Substances TSCA: Toxic Substances Control Act TCSI: Taiwan Chemical Substance Inventory INSQ: Inventario Nacional de Sustancias Químicas NCI: National Chemical Inventory FBEPH: Russian Register of Potentially Hazardous Chemical and Biological Substances

This document is copyright.

Apart from any fair dealing for the purposes of private study, research, review or criticism, as permitted under the Copyright Act, no part may be reproduced by any process without written permission from CHEMWATCH.

TEL (+61 3) 9572 4700.